PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISH	HED U	INDER THE PATENT COOPERATION TREATY (PCT)
(51) International Patent Classification 6:		(11) International Publication Number: WO 99/29788
C09D 11/00	A1	(43) International Publication Date: 17 June 1999 (17.06.99)
(21) International Application Number: PCT/GB (22) International Filing Date: 4 December 1998 (BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN,
(30) Priority Data: 9725928.7 5 December 1997 (05.12.97) (MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR,
(71) Applicant (for all designated States except US): XAA NOLOGY LIMITED [GB/GB]; Unit 316, Scien Cambridge CB4 0XR (GB).	R TEC nce Pa	H- GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF,
(72) Inventors; and (75) Inventors/Applicants (for US only): JOHNSON [GB/GB]; 3 Orchard Place, Hollygate, Helsby, WA6 OQE (GB). WOODS, Jill [GB/GB]; 10 V	Chesh	ire With international search report.

(54) Title: RADIATION CURABLE INK JET INK COMPOSITIONS

(74) Agents: CROPP, John, Anthony, David; Mathys & Squire, 100 Grays Inn Road, London WC1X 8AL (GB) et al.

Yaxley, Peterborough PE7 3LJ (GB).

(57) Abstract

A radiation curable ink jet ink comprising a colorant component, a diluent consisting essentially of reactive liquid material and, optionally, at least one photopolymerisation catalyst and wherein the reactive liquid material comprises monofunctional material, difunctional material and tri– or higher functional material and wherein the total amount of tri– or higher functional material in the ink forms more than 10 but not more than 30 % by weight of the total amount of reactive material in the ink, the total amount of monofunctional material in the ink is at least 20 % by weight of the total amount of reactive material in the ink, and the total amount of difunctional material in the ink is at least 17 1/2 % by weight of the total amount of reactive material in the ink and is such that the total amount of di– or higher functional material is not less than 35 % by weight, and the viscosity of the ink is not greater than 35 mPa.s at 30 °C.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AM AT AU AZ	Albania Armenia Austria Australia Azerbaijan	ES FI FR GA	Spain Finland France	LS LT LU	Lesotho Lithuania	SI SK	Slovenia Slovakia
AT AU AAZ	Austria Australia	FR			Limuania	O.M.	UIV TURIN
AU AZ	Australia		France		• • • • • • • • • • • • • • • • • • • •	SN	Senegal
AZ A		GA			Luxembourg		Swaziland
	A zerbaijan		Gabon	LV	Latvia	SZ	_
		GB	United Kingdom	MC	Monaco	TD	Chad
	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB 1	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Тигкеу
	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
-	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
	Belarus	IS	Iceland	MW	Malawi	US	United States of America
	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
	Central African Republic	JР	Japan	NE	Niger	VN	Viet Nam
	Congo	KE.	Kenya	NL	Netherlands	YU	Yugoslavia
	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
	••••	134	Republic of Korea	PL	Poland		
	Cameroon	KR	Republic of Korea	PT	Portugal		
-	China	KZ	Kazakstan	RO	Romania		
	Cuba	LC	Saint Lucia	RU	Russian Federation		
	Czech Republic	_	Liechtenstein	SD	Sudan		
	Germany	LI		SE	Sweden		
	Denmark	LK	Sri Lanka	SG	Singapore		
EE .	Estonia	LR	Liberia	36	Singapore		

RADIATION CURABLE INK JET INK COMPOSITIONS

This invention relates to ink jet ink compositions and in particular to such compositions which are curable by ultraviolet radiation.

Radiation curable inks e.g. for screen printing and offset printing have been known for some time and more recently, there have been proposals for radiation-curable ink jet inks; see, for example, GB-A-2233928, JP-A-63102936, US-A-4303924, EP-A-0540203, EP-A-0465039 and W097/31071. There are, however, substantial problems in arriving at a radiation-curable formulation which meets all the criteria required of an ink jet ink; specifically, low viscosity, the required level of surface tension, low volatility, long term stability, rapid drying of the ink dot derived from the droplet ejected from the nozzle of the print head, and the provision of print of satisfactory quality, especially resistance to scratching and adhesion to the substrate.

For example, the formulations described in JP-A-63102936 and US-A-4303924 include some unreactive diluent, such as ethyl alcohol, methyl alcohol, methylethylketone, ethyl acetate, or the like; however, including such a diluent presents the operator with the problem of removal of the diluent. Where a volatile diluent is used, this presents problems with stability of the ink during storage or in the

printer. In the compositions described in EP-A-0465039 and EP-A-0540203, on the other hand, which contain no unreactive diluent, the amount of polyunsaturated component that can be employed is limited because of its viscosity, thereby providing a constraint on the level of crosslinking that can be achieved on curing and thus on the properties of the print. For example, in the examples of EP-A-0465039, the functionality, that is the average number of unsaturated groups per molecule of the total of reactive components in the ink, is only about 1.22.

WO97/31071 proposes an ink jet ink composition for which a requirement is said to be that from 80 to 95%, based on the total weight of the composition, consists of alkoxylated and/or polyalkoxylated polyfunctional acrylate material. In our experience, however, compositions containing such levels of polyfunctional material, even if it is all alkoxylated or polyalkoxylated acrylate, tend to be too viscous for use in many applications.

We have now developed a radiation curable ink jet ink which is essentially free of non-reactive diluent and yet has the desired viscosity, surface tension, volatility, stability and drying rate and produces print of acceptable quality. Inks according to the invention thus can be printed without difficulty from ink jet printers to give good quality print on a variety of substrates including such disparate materials as paper and non-adsorbent materials e.g. metal.

They are of particular interest, however, in printing on plastics which are difficult to print on, e.g. polyolefins such as polyethylene and polypropylene.

According to one aspect of the present invention, there is provided a radiation curable ink jet ink comprising a colorant component, a diluent consisting essentially of reactive liquid material and, optionally, at least one photopolymerisation catalyst and wherein the reactive liquid material is formed of monofunctional material, difunctional material and tri- or higher functional material and wherein

the total amount of tri- or higher functional material in the ink forms more than 10 but not more than 30% by weight of the total amount of reactive material in the ink,

the total amount of monofunctional material in the ink is at least 20% by weight of the total amount of reactive material in the ink, and

the total amount of difunctional material in the ink is at least 17%% by weight of the total amount of reactive material in the ink and is such that the total amount of di- or higher functional material is not less than 35% by weight, and the viscosity of the ink is not greater than 35 mPa.s at 30°C as measured using a Brookfield Viscometer fitted with a small sample adaptor having a No. 18 spindle and at a rotational speed of 60 rpm, giving a shear rate of

79.2 sec⁻¹.

By consisting essentially of reactive liquid material, we mean that the diluent is free or substantially free of non-reactive components; that is to say, if any non-reactive liquid component is present in the diluent it forms no more than 1% by weight of the ink, preferably not more than 0.5%, by weight.

By reactive material we mean material containing one or more unsaturated carbon-carbon bonds polymerisable by radiation. The reactive material may comprise monomer or a mixture of monomer and oligomer. Oligomer is sometimes known as prepolymer. Non-limiting examples of monomers are acrylates, methacrylates and alkoxylated and polyalkoxylated derivatives thereof. Non-limiting examples of oligomers (prepolymers) are polyester-, urethane- and epoxy-acrylates.

In one preferred embodiment, the ink includes at least one polyfunctional oligomer. Preferably, said at least one oligomer has an equivalent weight of at least 180. Print obtained from inks containing such a component exhibits improved toughness, adhesion and/or scratch resistance as compared with print derived from the same ink composition but excluding the component.

Preferably, the oligomer is liquid so as to maintain the

desired low viscosity of the ink and preferably it has a functionality greater than 2. If it is not liquid, it should be soluble in the liquid components of the reactive material. In any event, the oligomer, if present, is deemed to form part of the reactive liquid material.

In another preferred embodiment, the ink composition includes a silicone derivative containing carbon-carbon unsaturation which is polymerisable by radiation, to adjust the surface tension of the ink. This polymerisable silicone derivative, if present, is also deemed to form part of the reactive liquid material.

While the possibility of formulating the compositions of the invention to be cured using other radiation sources, e.g. electron beam, is not excluded, the compositions are preferably formulated to be curable by visible, or more preferably ultra violet, light, in which case they will usually include at least one photoinitiator.

The components of the compositions of the invention will now be described in greater detail.

The colorant is preferably thermally stable and water-insoluble. While the possibility of using colorants, such as dyes, which are soluble in the diluent, is not excluded, it is preferred to use those, such as pigments, which are insoluble, especially where light-fastness is important.

In this case, it may be desirable to include a dispersant in the composition to stabilise the dispersion of insoluble colorant in the diluent.

Examples of insoluble colorants include, in particular, carbon black and those colorants characterised as pigment dyes in The Colour Index.

Examples of suitable pigments include those within the ranges having the following CI classifications:

Colour	CI Number
Green	PG 7 and 36
Orange	PO 5, 34, 36, 38, 43, 51, 60, 62,
	64, 66, 67 and 73
Red	PR 112, 149, 170, 178, 179, 185,
	187, 188, 207, 208, 214, 220,
	224, 242, 251, 254, 255, 260 and
	264
Magenta/Violet	PV 19, 23, 31, and 37 and PR 122,
	181 and 202
Yellow	PY 17, 120, 138, 155, 168, 175,
	179, 180, 181 and 185
Blue	PB 15
Black	PB 2, 5 and 7.

Examples of specific pigments include IRGALITE BLUE GLVO,

MONASTRAL BLUE FGX, IRGALITE BLUE GLSM, HELIOGEN BLUE L7101F, LUTETIA CYANINE ENJ, HELIOGEN BLUE L6700F, MONASTRAL GNXC, MONASTRAL GBX, MONASTRAL GLX, MONASTRAL 6Y, IRGAZIN DPP ORANGE RA, NOVAPERM ORANGE H5G70, NOVPERM ORANGE HL, MONOLITE ORANGE 2R, NOVAPERM RED HFG, HOSTAPERM ORANGE HGL, PALIOGEN ORANGE L2640, SICOFAST ORANGE 2953, IRGAZIN ORANGE 3GL, CHROMOPTHAL ORANGE GP, HOSTAPERM ORANGE GR, PV CARMINE HF4C, NOVAPERM RED F3RK 70, MONOLITE RED BR, IRGAZIN DPP RUBINE TR, IRGAZIN DPP SCARLET EK, RT-390-D SCARLET, RT-280-D RED, NOVAPERM RED HF4B, NOVAPERM RED HF3S, NOVAPERM RD HF2B, VYNAMON RED 3BFW, CHROMOPTHAL RED G, VYNAMON SCARLET 3Y, PALIOGEN RED L3585, NOVAPERM RED BL, PALIOGEN RED 3880 HD, HOSTAPERM P2GL, HOSTAPERM RED P3GL, HOSTAPERM RED E5B 02, SICOFAST RED L3550, SUNFAST MAGENTA 122, SUNFAST RED 122, SUNFAST VIOLET 19 228-0594, SUNFAST VIOLET 19 228-1220, CINQUASIA VIOLET RT-791-D, VIOLET R NRT-201-D, RED B NRT-796-D, VIOLET R RT-101-D, MONOLITE VIOLET 31, SUNFAST MAGENTA 22, MAGENTA RT-243-D, MAGENTA RT 355-D, RED B RT-195-D, CINQUASIA CARBERNET RT-385-D, MONOLITE VIOLET R, MICROSOL VIOLET R, CHROMOPTHAL VIOLET B, ORACET PINK RF, IRGALITE YELLOW 2GP, IRGALITE YELLOW WGP, PV FAST YELLOW HG, PV FAST YELLOW H3R, HOSTAPERM YELLOW H6G, PV FAST YELLOW, PALIOTOL YELLOW D1155 and IRGAZIN YELLOW 3R.

Mixtures of colorants may be employed, if desired, including mixtures of dyes, mixtures of pigments and mixtures of one or more dyes with one or more pigments.

In one preferred embodiment of the invention, the colorants are chosen to give the widest range of colours and tones in a hexachrome system.

The amount of colorant employed in the ink will depend on the choice of colorant and the depth of colour required in the print, and can be established by simple experiment. In general, for pigments it will fall within the range 0.01% to 50% by weight, the amount chosen being such that viscosity of the ink does not exceed 35 mPa.s. For organic pigments the amount will generally be in the range 0.01 to 10% weight, more preferably 0.05 to 6% most preferably 0.05 to 3%.

Where the colorant comprises a pigment which is to be dispersed in the diluent, it preferably has a particle size of not greater than $1\mu m$ maximum dimension and more preferably not greater than $0.5\mu m$. Even more preferably, the particles have a narrow size range distribution.

Where the ink comprises a dispersion of pigment, a dispersant will normally be required to aid or stabilise the dispersion. The choice of dispersant will depend upon the nature of the pigment and composition of the diluent. Examples of suitable materials may be found amongst dispersants sold under the trade names of Solsperse, EFKA and Byk. Mixtures of dispersants and mixtures of one or more dispersants with one or more dispersant synergists may

be employed. The amount of dispersant employed (or dispersant and synergist where used) will depend upon the choice and concentration of the pigment. For organic pigments, the amount will usually be in the range 15 to 100% by weight of the pigment, preferably 20 to 75% by weight. For inorganic pigments, lower concentrations may be acceptable, e.g. 5% or less.

The reactive liquid material of the ink composition comprises monofunctional, difunctional and tri- or higher By mono-, di-, tri- and higher functional material. meant compounds having, is material functional respectively, one, two, three or more unsaturated carboncarbon groups which are polymerisable by radiation, especially but not exclusively ultra-violet light. Examples of some suitable compounds may be found in the publication U.V. and E.B. Curing Formulations for Printing Inks, Coatings and Paints, edited by R. Holman and published by SITA-Technology, London, in 1984 and The Printing Ink Manual, Fourth Edition, published by VNR International, However, the choice of suitable compound is not limited to the compounds or classes of compounds disclosed in those publications. It is preferred to use compounds of low volatility, low viscosity and high reactivity. For the desired reactivity, therefore, it is generally preferred to employ compounds wherein the unsaturated carbon-carbon groups are vinyl CH_2 =CH-, with a lesser preference for vinylidene $CH_2=C$. Particularly

preferred are acrylates and methacrylates; however as methacrylates tend to be less active than the corresponding acrylates, they preferably form only a small proportion of the diluent where they are used.

The monofunctional material may comprise a single monomer or a mixture of monomers. Straight chain acrylates of lower alcohols (e.g. C_{1-4} alcohols) tend to be too volatile and therefore should form only a small proportion of the reactive liquid material or be avoided. Acrylates of branched chain alcohols e.g. isodecyl alcohol are less volatile but more preferred are acrylates of cyclic or polycyclic alkanols, e.g. tetrahydrofurfuryl acrylate and Examples of the monofunctional isobornyl acrylate. acrylates that may be used include long chain aliphatic acrylates e.g. wherein the aliphatic group contains at least 8, preferably at least 10 carbon atoms, such as lauryl acrylate and stearyl acrylate, and acrylates of alkoxylated alcohols e.g. 2-(2-ethoxyethoxy)ethyl acrylate. The monofunctional material need not necessarily be an acrylate; for example N-vinyl-2-pyrrolidone may be used. Further examples of monofunctional monomers are Sartomer CD 9050 (a monofunctional acid ester) and Sartomer CD 611 (an ethoxylated tetrahydrofuran acrylate).

So that the desired amounts of polyfunctional material may be incorporated into the ink, the overall viscosity of the monofunctional material of the diluent preferably is as low

as possible, e.g. 20 mPa.s or less, more preferably 16 mPa.s or less, at 30°C.

The tri- or higher functional material may also comprise a single monomer or a mixture of monomers. acrylates such as those obtained by acrylating the products of ethoxylating or propoxylating an initiator containing three or more active hydrogen atoms are particularly preferred. Examples of initiators having three or more include glycerol, trimethylol active hydrogen atoms propane, pentaerythritol and neopentyl alcohol. Examples of such alkoxylated acrylates are ethoxylated trimethylol propane triacrylates, propoxylated glyceryl triacrylates, Sartomer SR 9008 (an alkoxylated trifunctional acrylate ester) and propoxylated pentaerythritol tetraacrylates. Other examples of tri- or higher functional monomers are tris (2 - hydroxyethyl) isocyanurate triacrylate, (a trifunctional acrylate ester). Sartomer SR 9012 Preferably compounds having relatively low viscosities, e.g. 200 mPa.s or less at 30°C, are chosen so that amounts towards the upper end of the range of 10 to 30% by weight, may be employed in the diluent.

While compounds having a functionality higher than 3 may also be used, they should be included at most in only small amounts, the maximum tolerable amount being less as the functionality of the compound increases. This is because as the functionality increases, so does the crosslink

density in the cured product unless the molecular weight of the compound is increased but increase in molecular weight tends to lead to an increase in the viscosity of the compound.

The difunctional material, which may comprise a single monomer or mixture of monomers, makes up the balance of the reactive material. It is preferred that the viscosity of the difunctional material is not greater than 30mPa.s, and preferably not greater than 20 mPa.s, at 30°C. Preferred components are diacrylates of glycols and polyglycols, especially propylene glycol and polypropylene glycols e.g. di-, tri- and higher propylene glycols. Di- acrylates of tri- or higher hydric alcohols may also be used. Examples of such diacrylates having a viscosity at 30°C of not greater than 20 mPa.s are the diacrylates of 1,4butanediol, neopentylglycol, propoxylated neopentyl glycol, glycol, hexanediol, dipropylene glycol, diethylene tripropylene glycol, triethylene glycol and polyethylene glycols.

As indicated above, it is preferred that the reactive liquid material also includes at least one polyfunctional oligomer, so as to provide print of improved toughness, adhesion and/or scratch resistance. This oligomer may comprise any suitable compound or mixture of compounds containing at least two, and preferably more than two, unsaturated carbon-carbon bonds polymerisable by radiation

and an equivalent weight which is preferably at least 180, more preferably at least 200 or 220. However, in general, increase in equivalent weight is accompanied by increase in viscosity, thus restricting the amount of the oligomer that may be included in the composition. Preferably therefore, the equivalent weight does not exceed 750 and more preferably does not exceed 500.

As in the case of the other reactive components, oligomers which are polyacrylates are preferred because of their relatively low viscosity compared with other oligomeric polyfunctional materials. Examples include the polyfunctional products of acrylating hydroxy-terminated polyesters, known as polyester acrylates, the polyfunctional products of acrylating urethane oligomers, known simply as urethane acrylates, and epoxy acrylates. Alkoxylated acrylates are not regarded as oligomers.

It appears that there is an optimum concentration for the oligomeric material beyond which the improvement in properties is marginal. Since increasing the concentration tends to increase viscosity, it is generally not desirable to employ more than this optimum amount which will vary according to the choice of the oligomeric material and of the other components of the reactive liquid material. In general, this optimum will not exceed 30% of the reactive liquid material and will generally be not more than 25% e.g. 10 to 22%, more generally 15 to 20%, by weight of the

total amount of reactive liquid material in the ink. The amount of oligomeric material employed is to be included in the calculation of the overall amount of di- or tri- (or higher) functional material in the reactive liquid material, according to its functionality. Thus, if it is trifunctional, the total amount of oligomeric material and other tri- or higher functional material must be greater than 10% but not more than 30% by weight of the total amount of reactive liquid material in the ink.

The reactive liquid material should be essentially of low volatility and preferably substantially non-volatile at ambient temperatures and preferably also at the printhead temperature if above ambient.

A silicone derivative containing at least one radiation-polymerisable carbon-carbon unsaturation is a preferred component of the ink composition for reducing its surface tension, especially when the ink is intended for printing on plastics with low surface tension surfaces e.g. polyolefins and ABS. Preferably, it is employed in amount to give a surface tension below 35 dynes/cm and generally in the range 25 to 35 dynes/cm. In general, the silicone derivative will be used in amounts of 0.05-015 to 1-3% by weight, depending on the molecular weight of the derivative, and more preferably 0.1 to 0.6% by weight of the ink. The inclusion of the silicone derivative in such quantities also reduces the tendency to blocking of sheets

printed with the ink and improves dot definition. While greater amounts may further reduce surface tension, other properties are likely to be adversely affected. Examples of suitable compounds are silicone acrylates e.g. acrylates of organo modified silicones, for example organo modified polysiloxanes e.g. comprising repeating units of

hydrocarbyl group, e.g. methyl or phenyl, and at least one

unit of the structure - Si - O - where R is as defined X

above and X contains an acrylate (CH₂-CH.COO-) group. In one preferred embodiment, the acrylate group is connected to the silicon atom via an alkoxy or polyalkoxy group. Preferably the chosen silicone derivative will have one or two radiation-polymerisable unsaturated carbon-carbon bonds. While the silicone component is deemed to be part of the reactive liquid material of the composition, as the amount of the silicone derivative used in the composition is relatively small, it may be ignored when calculating relative the amounts of mono-, di- and/or tri- or higher functional material in the reactive liquid material.

The total amount of monofunctional material in the ink (including monofunctional oligomer if present) should form at least 20%, e.g. from 20 to 60% by weight, preferably from 20 to 50% by weight, and more preferably from 40 to

50% by weight of the total reactive material in the ink. If less than about 20% is employed, the viscosity of the ink will tend to be too high. If, on the other hand, the amount significantly exceeds 60%, the properties of the print, and in particular abrasion resistance and hardness may suffer, and also the rate and/or degree of cure of the ink may be reduced.

The total amount of tri- or higher functional material in the ink (including tri- or higher functional oligomer; if present) is greater than 10 but not more than 30% by weight of the total amount of reactive material in the ink. Preferably it is present in an amount of at least 15% by weight and more preferably from 20 to 30% by weight. If used in an amount of less than 10% by weight, the properties of the print obtained from the ink, and in particular hardness and scratch resistance, tend to suffer. On the other hand, if used in amounts of more than about 30% by weight, the viscosity of the ink tends to be too high and the print may become unacceptably brittle and inflexible.

The total amount of difunctional material (including difunctional oligomer, if present) is required to form at least 17% by weight of the total amount of reactive material in the ink and to be such that the viscosity of the ink is not greater than 35 mPa.s at 30°C and the diand higher functional material in the ink forms at least

35% of the total of reactive material in the ink. To achieve the desired combination of viscosity of the ink and flexibility of the print derived therefrom, preferably the difunctional material is present in an amount in the range 17% to 35% by weight of the total amount of reactive material in the ink, more preferably 20 to 30% by weight.

A feature that is relevant to the control of the properties of the print derived from the ink, and in particular its physical properties such as hardness, abrasion resistance, modulus, flexibility and elasticity, is the overall functionality of the reactive material in the ink; i.e. the average number of radiation polymerisable unsaturated carbon-carbon bonds per molecule. If this value is too low, the print will be too soft but if it is too high, the print will tend to be brittle and to craze or crack when the substrate on which it has been printed is non-absorbent, e.g. as in a plastic substrate, and is flexed. Preferably, the amounts of mono-, di- and tri- and higher functional components of the ink are selected such that this value will be in the range 1.25 to 2, more preferably 1.25 to 1.85, most preferably 1.3 to 1.7..

While inks according to the invention may be formulated for curing by any suitable form of electromagnetic radiation, visible light or, more preferably, ultra-violet light is preferred and for this purpose the ink will also preferably include at least one photoinitiator. Conventional

photoinitiators may be employed and the choice will depend upon the choice of colorant and the wave length of the radiation. Examples of suitable photoinitiators are:

- 2,2-dimethyl-2-hydroxy-acetophenone,
- 1-hydroxy-1-cyclohexyl-phenyl ketone,
- 2,2-dimethoxy-2-phenylacetophenone,
- 2-benzyl-2-dimethylamino-1-(4-morpholinophenyl)-butan-
- 1-one,
- 2-methyl-1-[4-(methylthio)phenyl]-2-morpholino-propan-
- 1-one,
- 2,4,6-trimethylbenzyl-diphenyl-phosphine oxide,
- 1-chloro-4-propoxythioxanthone,
- Isopropyl thioxanthone (mixture of 2- and 4- isomers),
 Benzophenone,
- Blends of bis (2,6-dimethoxybenzoyl)-2,4,4-trimethyl pentyl phosphine oxide and 1-phenyl-2-hydroxy-2-methyl propanone,
- Blends of bis (2,6-dimethoxybenzoyl)-2,4,4-trimethyl pentyl phosphine oxide and 1-hydroxy-cyclohexyl-phenyl ketone,
- bis (2,4,6-trimethylbenzoyl)phenylphosphine oxide, and Camphorquinone.

In order to obviate, so far as possible, the obscuring effect of the colorant, especially where this is a pigment, a mixture of photoinitiators is preferably used whose peak energy absorbtion levels are at different wave lengths

within the range of the selected radiation. In the inks of the present invention, preferential absorption of incident UV radiation by the pigments leads to depletion of the available energy for the photoinitiator(s). This can place considerable constraints on the cure speed and level of conversion achievable especially with black pigmented ink formulations. Ιt is therefore preferred to employ photoinitiator blends which are sensitive to the wavelengths not absorbed, or only partially affected, by the pigments. Typically this "absorption window" between 350-400nm.

The initiators will generally be used in amounts of from about 3 to about 15% more usually about 5 to about 10% by weight of the ink.

Activators, e.g. co-initiators or amine synergists, for the photoinitiators may also be included, if desired. Examples include ethyl-4-(dimethylamino) benzoate, 2-ethylhexyl dimethylaminobenzoate, and dimethylaminoethyl methacrylate.

In operation, the ink is ejected from an ink jet printer and exposed to the selected radiation during its flight from the printhead to the substrate to be printed or, more preferably after deposition on the substrate. Generally speaking, the smaller the throw distance, that is the distance from the printhead to the substrate, the better the print quality. However it is preferably no less than

0.5 mm and even at distances of 1.5 mm or 2 mm or even higher, acceptable results may be obtained. For most applications, the preferred throw distance will be in the range 0.5 to 0.75 mm. Curing of the ink is aided by the absence of oxygen and thus it is preferred to provide an inert gas atmosphere, e.g. nitrogen over the substrate in the zone where the ink is exposed to the radiation. Curing is preferably effected using radiation in the UVA and near-visible frequencies.

Inks according to the invention may be formulated for use in any of the available kinds of continuous and drop-ondemand ink jet printers although in general they may not be suitable for use in apparatus which employs thermal means of droplet generation. In a continuous ink jet printer, for example, former kind, for example, ink droplets which are produced continuously may be passed through a charging area where individual droplets receive an electrical charge in response to a signal and are directed towards a substrate to be printed. The droplets then pass through an electrical field causing them to be deflected by an amount which is dependent on the intensity of the charge and the Droplets not required to form print on the substrate may be directed to a by-pass gutter. For inks of the present invention to be suitable for use in such printers, they are preferably conductive and it may therefore be necessary to include a suitable conducting additive. Drop-on-demand ink jet printers may be of the

kind using an electrostatically accelerated ink jet or droplet sequences ejected by pressure impulse actuation, e.g. when each droplet is individually ejected from a nozzle by means of pressure pulses induced e.g. by means of a piezoelectric actuator acting on the ink in the channel supplying the nozzle.

The diluent, colorant(s), dispersant(s) (where present), dispersant synergist(s) (where present), photoinitiator(s) including activator(s) therefor (where present), and conducting additive (where present) will normally form at least 99% of the composition and preferably substantially the entire composition. As indicated above, preferably at least 99.5% by weight of the diluent is reactive liquid material.

The inks may be employed for printing on to a wide variety of substrates, both absorbent and non-absorbent including paper, glass, plastic and metal, e.g. steel, copper and aluminium, but are particularly suitable for printing on to plastics to provide a strongly bonded print of good definition and optical density, especially if the surface to be printed is pre-treated e.g. by flame, plasma etch or corona treatment to raise the surface energy. Preferably the surface energy should be at least about 36 dynes/cm and more preferably from 42 to 48 dynes/cm.

Examples of plastics on which the inks of the present

invention have been successfully printed are polyolefins such as polyethylene (including high density polyethylene), and polypropylene, vinyl chloride polymers, ABS and foamed plastics such as expanded polystyrene.

The invention is now illustrated but in no way limited by the following Examples, in which all parts are expressed as parts by weight except where otherwise indicated.

Example 1

The compositions set out in Table 1 below were formulated into inks in the manner now described.

Preparation of Millbase

The pigment was ground, with the appropriate amount of hyperdispersant to ensure minimum millbase viscosity, using conventional bead milling techniques. The millbase so formed was processed until a fine particle dispersion was obtained. The particle grind was assessed by visual microscopy and a filterability test. The finished millbase was removed from the grinding media using an appropriately sized mesh either by pressure or gravity.

Preparation of Ink from Millbase

All the following blending operations were carried out in

opaque vessels (either amber glass or stainless steel) using an electrically operated high speed mixer fitted with a stainless steel rotor blade. A homogenous mixture was formed of the reactive diluent(s) and oligomers. The photoinitiator(s), either in solid or liquid form, were then added and stirring/heating continued until all insoluble components dissolved. The millbase was then carefully added to the stirred/heated vehicle in a manner to avoid any unwanted 'shock' crystal seeding/growth. Finally, the silicone polyether acrylate was added and stirring/heating continued until visual homogeneity was achieved.

The ink composition so formed was filtered using a proprietary cartridge filter rated at 1 micron absolute filter rating. Following bottling a retained sample was analysed for viscosity, visual appearance, cure rate, surface tension and ageing at elevated temperature (60°C).

The viscosities of the ink compositions were all in the range 15-20cps measured using a Brookfield Viscometer at 30°C and their values for surface tension were all in the range of 26-28 dynes/cm.

The inks were used to form coloured print on moulded ABS cards using a 128 channel greyscale drop-on-demand printhead of the kind described for example in EP-A-0.277.703 and EP-A-0.278.590 with a nozzle diameter of

 $25\mu\text{m}$ and a distance of 2mm between the printhead and the card. Curing was by means of a Fusion F300s UV Lamp with a 152mm (6 inch) long "D bulb" at 11.8 w/mm (300w/inch) using cure energies in the range 1-3J/cm². Curing was complete within one second.

The print was well defined with acceptable colour density. The cyan, magenta and black inks all exhibited colour densities greater than 1.5 and were typically about 1.8. The colour density of the yellow ink was greater than 1, typically 1.1.

The print exhibited good adhesion to the card and good scratch resistance. Scratching with a fingernail did not remove the print and on further testing of adhesion by scoring and crosslinking the print with a scalpel and attempting to remove it by applying Sellotape to the scored surface and then peeling off the Sellotape, no detectable removal of ink was observed. The hardness of the print was assessed in conventional manner using pencils of different hardnesses. The hardness is determined as the lowest hardness value to form a scratch on the print. Print formed from the inks of the present invention generally exhibited hardnesses of 6H-7H.

The print remained unaffected when the card was subjected to the dynamic bending stress test in accordance with International Standard ISO/IEC 10 372.

TABLE 1

	BLACK	CYAN	MAGENTA	YELLOW	CYAN	MAGENTA
ACTILANE 430	10%	10.00%	10%	10%	10%	109
ACTILANE 251	15%	15.00%	15%	15%	15%	15%
TEGORAD 2200	0.40%	0.40%	0.40%	0.40%	0.40%	0.40%
ISOBORNYL ACRYLATE	39.70%	40.00%	38.35%	41.48%	40.85%	41.47%
SARTOMER 306	23.00%	22.90%	23.00%	22.0%	23.67%	23.00%
SPEEDCURE ITX	2.00%	-	-	_	-	_
QUANTACURE EHA	3%	-	_	_	-	_
IRGACURE 907	5.00%	-	-	-	-	_
REGAL 250R	1.50%	-	_	_	-	_
SOLSPERSE 24000	0.38%	0.60%	0.75%	0.30%	0.03%	0.03%
SOLSPERSE 5000	0.03%	0.11%	_	-	0.01%	_
IRGALITE BLUE GLVO	-	1.00%	-	-	0.05%	_
HOSTAPERM RED	•	-	2.50%	_	_	0.10%
PALIOTOL YELLOW		-	•	0.75%	-	-
SOLSPERSE 22000		-	-	0.07%	-	_
LUCERIN TPO	-	5.00%	5.00%	5.00%	5.00%	5.00%
DAROCURE 1173	_	5.00%	5.00%	5.00%	5.00%	5.00%
SPEEDCURE EDB	-		-	_	-	
IRGACURE 369	_	-	_	_		

KEY:

Actilane 430 - trimethylpropane ethoxylate triacrylate (ex Akcros)

Actilane 251 - trifunctional urethane acrylate prepolymer (ex Akcros)

TegoRad 2200 - silicone polyether acrylate (ex Tego Chemie Service)

Isobornyl acrylate (Ex Cray Valley Products)

Sartomer 306 - tripropylene glycol diacrylate (ex Cray Valley Products)

Speedcure ITX - isopropylthioxanthone (ex Lambsons)

Speedcure EDB - ethyl 4-(dimethylamino)benzoate (ex Lambsons)

Irgacure 369 -2-benzyl-2-diethylamino-1-(4morpholinophenyl)-butanone-1

Solsperse 5000/22000/24000 - hyperdispersants (ex Zeneca) Regal 250R - carbon black, C.I. pigment black 7 (ex Cabot) Irgalite Blue GLVO - copper phthalocyanine, C.I, pigment blue 15:4 - (ex Ciba Geigy)

Hostaperm Red E5B 02 - quinacridone, C.I. pigment violet 19
- (ex Hoechst)

Paliotol Yellow - isoindoline, C.I. pigment yellow 185 - (ex BASF)

Quantacure EHA -2-ethylhexyl p-dimethylaminobenzoate (ex Great Lakes)

Irgacure 907 - 2-methyl-1-(4-methylthio)phenyl-2-morpholino-propan-1-one (ex Ciba Geigy)

Darocure 1173 - 1-phenyl-2-hydroxy-2-methylpropane (ex Merck)

Lucirin TPO - 2,4,6-trimethylbenzoyl-diphenyl-phosphine oxide (ex BASF).

The functionality of the reactive material of each of the above formulations was approximately 1.5.

Example 2

In the same manner as Example 1, a further series of ink jet ink formulations were prepared having the compositions set out in Table 2 below. Coloured print was formed from the inks in the manner described in Example 1 and similar results were obtained.

TABLE 2

	ORANGE	VIOLET	GREEN
ACTILANE 430	10%	9.55%	10%
ACTILANE 251	15%	14.32%	15%
TEGORAD 2200	0.40%	0.38%	0.40%
ISOBORNYL ACRYLATE	39%	21.41%	38.8%
N-VINYL PYRROLIDONE	_	19.09%	-
SARTOMER 306	23%	23.37%	23%
SPEEDCURE ITX		_	-
QUANTACURE EHA		-	-
IRGACURE 907		-	-
REGAL 250R	-	-	-
SOLSPERSE 24000	0.6%	0.535%	0.64%
SOLSPERSE 5000	_	0.135%	0.16%
CROMOPHTHAL ORANGE GP	2%	<u>-</u>	-
CROMOPHTHAL VIOLET B	_	1.67%	
MONASTRAL GNX-X	-	_	2%
SOLSPERSE 22000	-		_
LUCERIN TPO	5%	4.77%	5%
DAROCURE 1173	5%	4.77%	5%
SPEEDCURE EDB	_		-
IRGACURE 369		<u>-</u>	_

Example 3

In a series of further experiments it was found that incrementally replacing the Actilane 251 by Actilane 430 reduced the scratch resistance of the resultant cured print when tested following ASTM test method D 3363-92a, although merely increasing the Actilane 430 to 14%, ie. about 16% by weight of the reactive material, while reducing the Actilane 251 to 11% made no apparent difference to the

scratch resistance. If the amount of Actilane 251 is increased to 25% or 30% at the expense of Actilane 430 and Sartomer 306 the viscosity tends to increase without a sufficiently concomitant beneficial increase in physical properties.

Example 4

In the same manner as described in Example 1, a further series of ink jet ink formulations were prepared having the compositions set out in Table 3.

Coloured print was formed from the inks in the manner described in Example 1 and similar results were obtained.

TABLE 3

	BLACK	CYAN	MAGENTA	AETFOM
ACTILANE 430	12.50%	12.50%	12.50%	12.50%
ACTILANE 251	12.50%	12.50%	12.50%	12.50%
TEGORAD 2200	0.40%	0.40%	0.40%	0.40%
SARTOMER 506*	39.70%	40.00%	38.35%	41.48%
SARTOMER 306	23.00%	22.90%	23.00%	22.0%
SPEEDCURE ITX	2.00%	-	-	-
QUANTACURE EHA	3.00%	_		_
IRGACURE 907	5.00%	_	-	-
REGAL 250R	1.50%	-	-	_
SOLSPERSE 24000	0.38%	0.60%	0.75%	0.30%
SOLSPERSE 5000	0.03%	0.11%		-
IRGALITE BLUE GLVO	-	1.00%	-	_
HOSTAPERM RED E5B 02	-	-	2.50%	-
PALIOTOL YELLOW D1155	-	-	-	0.75%
SOLSPERSE 22000		-	-	0.07%
LUCERIN TPO	-	5.00%	5.00%	5.00%
DAROCURE 1173	_	5.00%	5.00%	5.00%

* Sartomer 506 is isobornyl acrylate (ex Cray Valley Products)

Example 5

A white ink was prepared having the following formulation:

Actilane 422	18.55%
Actilane 251	15%
N-vinyl pyrrolidone	20%
Solsperse 24000	1.05%
Tegorad 2200	0.4%
Irgacure 907	10%
White pigment	35%

Actilane 422 is dipropylene glycol diacrylate (ex Akcros). The white pigment is Tioxide TR92.

White print was formed from the ink in the manner described in Example 1 and similar results were obtained.

CLAIMS:

1. A radiation curable ink jet ink comprising a colorant component, a diluent consisting essentially of reactive liquid material and, optionally, at least one photopolymerisation catalyst and wherein the reactive liquid material comprises monofunctional material, difunctional material and tri- or higher functional material and wherein

the total amount of tri- or higher functional material in the ink forms more than 10 but not more than 30% by weight of the total amount of reactive material in the ink,

the total amount of monofunctional material in the ink is at least 20% by weight of the total amount of reactive material in the ink, and

the total amount of difunctional material in the ink is at least 17% by weight of the total amount of reactive material in the ink and is such that the total amount of di- or higher functional material is not less than 35% by weight, and the viscosity of the ink is not greater than 35 mPa.s at 30°C as measured using a Brookfield Viscometer fitted with a small sample adaptor having a No. 18 spindle and a rotational speed of 60 rpm, giving a shear rate of 79.2 sec-1.

- 2. An ink jet ink as claimed in claim 1 characterised in that the reactive liquid material includes at least one polyfunctional oligomer.
- 3. An ink jet ink as claimed in claim 2 characterised in that said at least one polyfunctional oligomer has a functionality greater

than 2.

4. An ink jet ink as claimed in claim 2 or claim 3 characterised in that said at least one polyfunctional oligomer is selected from polyester acrylates, urethane acrylates and epoxy acrylates.

- 5. An ink jet ink as claimed in any one of claims 2 to 4 characterised in that said at least one polyfunctional oligomer is present in the ink in an amount of up to 25% by weight of the total amount of reactive material in the ink.
- 6. An ink jet ink as claimed in any one of claims 2 to 4 characterised in that said at least one polyfunctional oligomer is present in the ink in an amount of from 10 to 22% by weight of the total amount of reactive material in the ink.
- 7. An ink jet ink as claimed in any one of claims 1 to 6 further including a silicone derivative containing carbon-carbon unsaturation which is polymerisable by radiation.
- 8. An ink jet ink as claimed in claim 7 characterised in that the silicone derivative is selected from silicone acrylates.
- 9. An ink jet ink as claimed in claim 7 or claim 8 characterised in that the silicone derivative contains repeating units have the structure

and at least one unit the having the structure

where each R is monovalent hydrocarbyl and X contains an acrylate group.

- 10. An ink jet ink as claimed in claim 5 characterised in that R is methyl.
- 11. An ink jet ink as claimed in any one of claims 7 to 10 characterised in that the silicone derivative is employed in an amount such that the ink has a surface tension of 25 to 35 dynes/cm.
- 12. An ink jet ink as claimed in any one of claims 7 to 11 characterised in that the silicone derivative forms from 0.05 to 1% by weight of the ink.
- 13. An ink jet ink as claimed in any one of the preceding claims characterised in that the colorant comprises a pigment dispersed in the reactive diluent.
- 14. An ink jet ink as claimed in claim 13 further including a dispersant for the pigment.

15. An ink jet ink as claimed in any one of the preceding claims characterised in that the colorant forms from 0.01 to 50% by weight of the ink.

- 16. An ink jet ink as claimed in any one of the preceding claims characterised in that the colorant forms from 0.1 to 10% by weight of the ink.
- 17. An ink jet ink as claimed in any one of the preceding claims characterised in that the components of the reactive liquid material are selected from acrylates, methacrylates and N-vinyl pyrrolidone.
- 18. An ink jet ink as claimed in any one of the preceding claims characterised in that the viscosity of the monofunctional material of the reactive diluent is not greater than 20 mPa.s at 30°C.
- 19. An ink jet ink as claimed in any one of the preceding claims characterised in that the monofunctional material of the diluent comprises at least one monomer selected from long chain alighatic acrylates, acrylates of branched chain alcohols, acrylates of alkoxylated alcohols, acrylates of cyclic and polycyclic alkanols, and N-vinyl pyrrolidone.
- 20. An ink jet ink as claimed in claim 18 or claim 19 in which the monofunctional material comprises isobornyl acrylate.
- 21. An ink jet ink as claimed in any one of the preceding claims characterised in that the tri- or higher functional material of the

reactive diluent comprises at least one monomer selected from ethoxylated trimethylol propane acrylates and propoxylated glyceryl acrylates.

- 22. An ink jet ink as claimed in any one of the preceding claims characterised in that the difunctional material of the reactive diluent has a viscosity of not greater than 20 mPa.s at 30°C.
- 23. An ink jet ink as claimed in any one of the preceding claims characterised in that the diffunctional material of the reactive diluent comprises at least one monomer selected from diacrylates of glycols and polyglycols.
- 24. An ink jet ink as claimed in any one of the preceding claims characterised in that the diffunctional material of the reactive diluent comprises at least one monomer selected from the diacrylates of 1,4-butanediol, neopentyl glycol, propoxylated neopentyl glycol, hexanediol, dipropylene glycol, tripropylene glycol, diethylene glycol, triethylene glycol and polyethylene glycols.
- 25. An ink jet ink as claimed in any one of the preceding claims characterised in that the monofunctional material forms 20 to 60% by weight of the reactive liquid material.
- 26. An ink jet ink as claimed in claim 25 characterised in that monofunctional material forms not more than 50% by weight of the reactive liquid material.

27. An ink jet ink as claimed in any one of the preceding claims characterised in that the tri- or higher functional material forms from 20 to 30% by weight of the reactive liquid material.

- 28. An ink jet ink as claimed in any one of the preceding claims characterised in that the difunctional material forms at least 20% by weight of the reactive liquid material.
- 29. An ink jet ink as claimed in any one of the preceding claims characterised in that the difunctional material forms not more than 35% by weight of reactive liquid material.
- 30. An ink jet ink as claimed in any one of the preceding claims characterised in that the overall functionality of the reactive material in the ink is in the range 1.25 to 2.
- 31. An ink jet ink as claimed in any one of the preceding claims characterised in that the overall functionality of the reactive material in the ink is in the range 1.25 to 1.85.
- 32. An ink jet ink as claimed in any one of the preceding claims characterised in that the overall functionality of the reactive material in the ink is in the range 1.3 to 1.7.
- 33. An ink jet ink as claimed in any one of the preceding claims characterised in that it includes at least one photoinitiator and is curable by ultra-violet light.

34. An ink jet ink as claimed in claim 33 characterised in that it includes two or more photoinitiators.

- 35. An ink jet ink as claimed in claim 33 or 34 characterised in that said at least one photoinitiator promotes cure in the frequency range 350-400nm.
- 36. An ink jet ink as claimed in any one of claims 33 to 35 characterised in that said at least one photo initiator forms from 3 to 15% by weight of the ink.
- 37. An ink jet ink as claimed in any one of claims 33 to 35 characterised in that said at least one photo initiator forms from 5 to 10% by weight of the ink.
- 38. An ink jet ink as claimed in any one of the previous claims comprising diluent, colorant, and optionally one or more of dispersant, dispersant synergist, photoinitiator, activator for photoinitiator and conducting additive and wherein said diluent, colorant, dispersant, dispersant synergist, photoinitiator, activator for photoinitiator and conducting additive form together at least 99% by weight of the composition.
- 39. A method of printing on a substrate comprising directing an ink as claimed in any one of claims 1 to 38 from the printhead of an ink jet printer towards a print-receiving surface of said substrate and, during its travel from the printhead to the substrate and/or after deposition on the substrate, exposing the ink to radiation to cure

the ink.

40. A method as claimed in claim 39 characterised in that the ink jet printer is a drop-on-demand printer.

- 41. A method as claimed in claim 39 or claim 40 characterised in that the surface is plastic.
- 42. A method as claimed in claim 41 characterised in that the surface is ABS or polyolefin.
- 43. A method as claimed in claim 41 or claim 42 characterised in that the ink has a surface tension of 25-35 dynes/cm.
- 44. A method as claimed in any one of claims 41 to 43 characterised in that the surface is pre-treated by flame or corona treatment.
- 45. A method as claimed in any one of claims 39 to 44 characterised in that the distance from the printhead to the substrate is in the range 0.5 to 1.5 mm.
- 46. A method as claimed in any one of claims 39 to 45 characterised in that the ink is exposed to the radiation in an inert atmosphere.
- 47. A printed substrate having a surface carrying print comprising a cured ink composition as claimed in any one of claims 1 to 38.
- 48. A printed substrate as claimed in 47 characterised in that the

surface is plastic.

49. A printed substrate as claimed in 48 characterised in that the surface comprises ABS or polyolefin.

INTERNATIONAL SEARCH REPORT

Inter: mai Application No

· ·		PC1/GB	98/03628
A. CLASS IPC 6	SIFICATION OF SUBJECT MATTER C09D11/00		***
L .	to International Patent Classification (IPC) or to both national classif	ication and IPC	
	S SEARCHED ocumentation searched (classification system followed by classification system followed by classi	Alexandra Land	
IPC 6	CO9D CO8D	tion symbols)	
Documenta	tion searched other than minimum documentation to the extent that	such documents are included in the field	ds searched
Electronic o	data base consulted during the international search (name of data b	ase and, where practical, search terms (used)
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the re	November 1	<u> </u>
	whole appropriate, of the re-	nevant passages	Relevant to claim No.
X	WO 97 31071 A (COATES BROTHERS P NIGEL ANTONY (GB); SELMAN HARTLE 28 August 1997	LC ;CAIGER Y DAVI)	1
	cited in the application see claims 1-14; examples 1,5		
X	EP 0 555 069 A (SERICOL LTD) 11 August 1993 see claims 1-14; example 1		1
Α	EP 0 465 039 A (DOMINO PRINTING PLC) 8 January 1992 cited in the application	SCIENCES	
A	EP 0 540 203 A (DOMINO PRINTING PLC) 5 May 1993	SCIENCES	
	cited in the application		
<u> </u>	er documents are listed in the continuation of box C.	Patent family members are list	ed in annex.
	egories of cited documents :	"T" later document published after the	nternational filing date
COUSIDE	nt defining the general state of the art which is not ared to be of particular relevance ocument but published on or after the international	or priority date and not in conflict w cited to understand the principle or invention	theory underlying the
"L" documer	ate	"X" document of particular relevance; the cannot be considered novel or can	not be considered to
citation	s cred to establish the publication date of another or other special reason (as specified)	involve an inventive step when the "Y" document of particular relevance; the cannot be considered to involve an	e claimed invention
otherm		ments, such combined with one or ments, such combination being ob-	more other such docu-
"P" documer later tha	nt published prior to the international filing date but an the priority date claimed	in the art. "&" document member of the same pate	
Date of the a	ctual completion of the international search	Date of mailing of the international	
3	February 1999	11/02/1999	
Name and m	ailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2	Authorized officer	
	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Stienon, P	

INTERNATIONAL SEARCH REPORT

information on patent family members

Interr nal Application No
PCT/GB 98/03628

			T T	·		90/03028
	atent document d in search repor	rt	Publication- date		Patent family member(s)	Publication date
WO	9731071	Α	28-08-1997	AU EP	1804597 A 0882104 A	10-09-1997 09-12-1998
EP		Α	11-08-1993	GB DE DE ES JP US	2264118 A 69300984 D 69300984 T 2082588 T 6001809 A 5395863 A	18-08-1993 25-01-1996 14-08-1996 16-03-1996 11-01-1994 07-03-1995
EP	0465039	Α	08-01-1992	DE DE JP US	69106952 D 69106952 T 5214279 A 5275646 A	09-03-1995 24-05-1995 24-08-1993 04-01-1994
EP	0540203	Α	05-05-1993	DE DE JP US	69215835 D 69215835 T 5214280 A 5275646 A	23-01-1997 03-04-1997 24-08-1993 04-01-1994